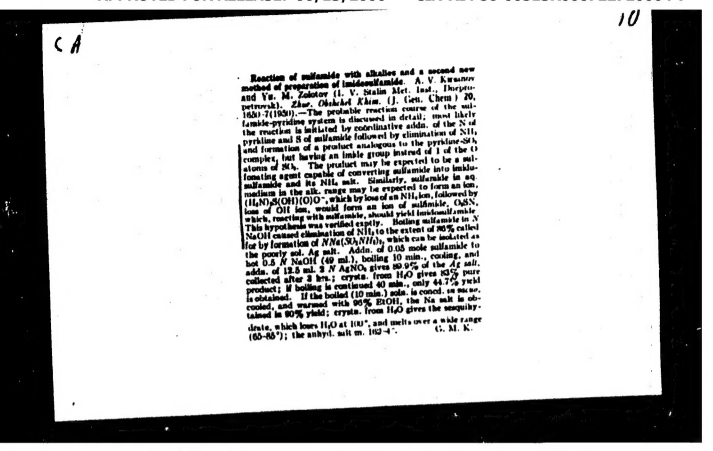


CA

Mechanism of amidation by means of sulfamide and a new method of preparation of imideoulfamide. A. V. Kiranov and Yu. M. Zolutov (I. V. Stalin Met. Inst., Theopropetrovak). Zhar. Obshchel Khim. (J. Gen. Chem.) 20, 1627-40(1920).—In the amidation of RCOsH by (HsN):-30, in pyrifine, the latter is not the only amidating agent, as the NH, mit of imideoulfamide (cf. C.A. 4, 426) that forms in the action of pyrifine on sulfamide also takes part. No stable intermediates form in the reaction and RCOsH; form as such in the mist. Heating 1.22 g. B20H with 0.9g. sOs(NH₀); and 5 mi. pyrifine 3 hrs. to 100°, evapn. st sector at 80-100°, evain. with also HtQ, and evapn. of the ext. gave 1.1 g. evain. with also HtQ, and evapn. of the ext. gave 1.1 g. evain. with also HtQ, such and 0.22 g. BrOH. Heating 4.8 g. SO₂(NH₀); with 10 ml. dry pyridine gave in 3 hrs. a viscous, insol. oll, which, extd. with MecCO, gave 9.37% (H₁N);SO₂ and 4.4 g. (91.7%) sirup, having the same compon. as sulfamide, sol. in HtQ, neutral, insol. in Et₂C and Me₂CO. If traces of H₂O are present, NH₂ is evalved during the reaction; in the presence of an equiv. amt. of H₂O a theoretical amt. of NH₃ is expelled. The sirup forms in 30 min. if the reaction is run at 130°. The behavior of the sirup indicates its structure is that of the NH₄ sait of imideoulfamide, N(N/H₂)(SO₂N/H₂). Heating it with p-O₃NC₄H₂Co₄H on a water lath readily gave p-O₃NC₄H₂Co₄H on a water lath readily gave p-O₃NC₄H₂Co₄H on a large with H₂O causes development of acality and evolution of NH₃, leaving behind

sulfamide. The aq. soln. of the strup with 2 N AqNO₃ yields cryst. NAq(SO₃NH₂), 2H₂O₃ softening at 100°, m. 105-7°; the meanm-decol product m. 108-9°; recrystn. from hot H₂O gave servated crystals from concel, soln.; the product is to 1:1150 in H₂O at 20°. Possibly the solt is chelated by hydrogen-bonding across the 2 terminal NH₂ groups and by Aq forming a 2nd bridge in same positions. The NH₃ solt lones 0.25 of its N very rapidly on boding in aq. soln., followed by slower formation of H₂NSO₄N₃.

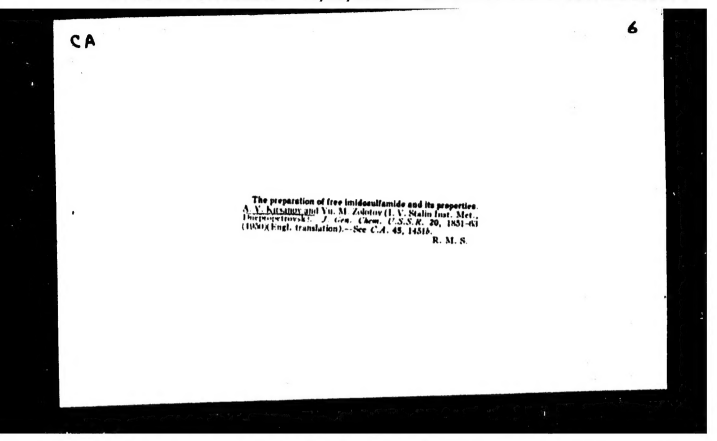
G. M. Knaolapoff

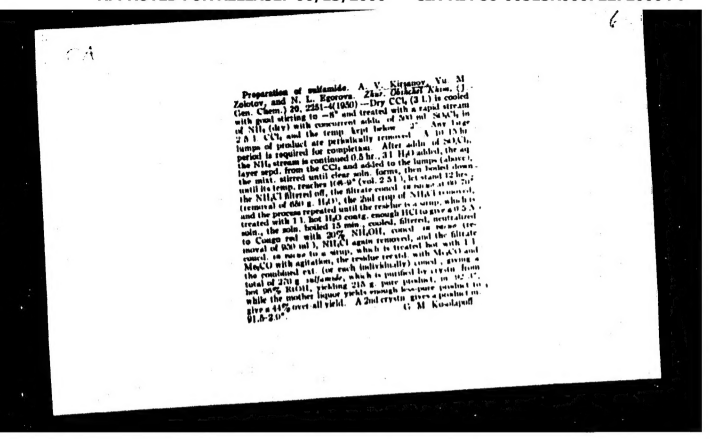


CA

Proc imidentifamide and its properties. A. V. Kirsanov and Yu. M. Zolotov (I. V. Stalin Metallurg. Thm., Deepto-petrovsk). Zhur. (bricket Khim. (I. Cen. Chem.) 20, 1790-1801(1960).—The free NH₂SO_NHBO_NMH₂ (I) was 1790-1801(1960).—The free NH₂SO_NHBO_NMH₃ (I) was obtained in 80% yield by decompa, of the Na salt in aquola, with 0.9 equiv. of H₂SO_N at 0° and subsequent evaps, at 0°. Yields up to 90% with respect to the recrystd. Na salt were obtained with the use of a mixt. of 90% of the choweteria amt. of H₂SO_N with a slight excess of NH₂SO_M. The crude I m. 164-70°; once-recrystd. material m. 162-4°, twice-recrystd. m. 167-9°; no further change in m.p. occurred on subsequent recrystas. The product is readily and, with evolution of heat, in H₂O and Me₇CO, sol. in McCOIII and is KtOA-, practically insel, in CdI₂. The contents of in 100 g. of said, sain, at 20° are: in H₂O, 30.1-30.4 g., in Me₇CO 33.0-35.9 g., in McCOC CdI₃ 1:3, a shout 0.64 g. The best method of recrysta, is soln, in dry 810Ac at room temp., filtration, and evaps. to drysess in sacros at room temp., soln, of the residue in a small ant, of about 0.64 g. The best method of recrysta, is soln, in dry 1810Ac at room temp., soln, of the residue in a small ant, of about 1810O, H., which is insol, in KtOAc but is sol, in Me₇CO to the extent of 0.3 g./100 g. After a 2nd recrystantered with EtOAc is necessary for the removal of the N11, SO, H., which is insol, in KtOAc but is sol, in Me₇CO to the extent of 0.3 g./100 g. After a 2nd recrystantered with EtOAc sides evaps. 1-2° lower than the fine needles pptd. from these solns, by CdI₄. The compen corresponds to the formula H₁, N₂SO₃, but cryoscopic deta, in 11₂O gives a 1101, wt.

roughly nail that formula. Complete hydrolysis gives I + II O = Nif SO, Nif + Nif SO, II, which confirms the stream of formula. I is a strong acts and is accurately streated both with phenolphthalein and with methyl orange, thrated both with phenolphthalein and with methyl orange, By colorimetry, a 0.01 N sola. 3.05. By cryscopy, 0.01-0.03 N sola. of 0.001 N sola. 3.05. By cryscopy, 0.01-0.03 N sola. of 1 is HaO are almost completely dissord. The rate of hydrolysis of I in HaO could be followed by analysis based on the difference of soly, between I and Nil SO, II (a product of the hydrolysis in EtOAc. The lat-order rate const. R of the hydrolysis in 0.25 N sola., at 29, 40, and 30°, was dettl. to 10% = 0.111, 1.78, and 6.17, resp., and the half-times to 17.3 hrs., 65, and 19 min., resp. These data are accurate only to within \$5%. They definitely contradict accurate only to within \$5%. They definitely contradict Hantasch's (Ber. 34, 343)(1901)) assertion of instantaneous hydrolysis of I in II/O. The following salts of I were prepared (NIIBO), N.Nils, sol. in II/O, sparingly wil, in cold, somewhat more is warm EtOH, transparent thin sandles, son-hygroscopic, softening at 114°, m. 115-18° to a turbid liquid; the sq. sola. is neutral to Congo. The Ag salt liquid; the sq. sola. is neutral to Congo. The Ag salt sold is klentical with that perpel, directly equiv. amt. of Ag.NO, is hientical with that perpel, from 1 and NaOH on SO,(Nils). The Ba salt, Ba(NilsO, sol., sol., naIIO, sparingly sol. in EtOH, decomp at 228-33°. The Hg and the Cu salts, not further levestigated, decomp. 31-13°. Call, N.H., N.S.O., sol. in II/O, sparingly sol. in cold RtOH and Me₂CO, softening at 130°, and solid liquid becoming the 133°. The solid liquid becoming clear at 135°. The solid liquid becom





KIPSANOV, A. V.

USER/Chemistry - Organic Chemistry

Apr 51

"Methylation of Imidosulfamide," A. V. Kirsanov, Tu. M. Zolotov, Chair of Org Chem, Dnepropetrovsk Metallurgical Inst

"Zhur Obshch Khim" Vol XXI, No 4, pp 642-645

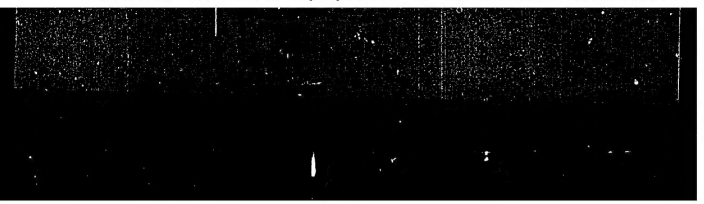
Prepd 3-monomethylimidosulfamide. Describes its properties. Notes that methylimidosulfamide reacts with NH3 in either soln to yield binary compd with curious properties, to be described in future report.

182716

KIRSANCV, A. V.

"The dimethylamidation of carboxylic acids." A. V. Kirsanov and Yu. M. Zolotov. (p. 1166)

SO: Journal of General Chemistry (Zhurnal Obshchei Khimii) 1951, Vol 21, No 6.



KIRSANOV, A. V.

Card 1 of 2

USSR/Chemistry - Phosphorus-Sulfur- Jul/Aug 52. Witrogen Compounds

"New Data on the Chemistry of Sulfuric Acid Amides and Trichlorophosphazosulforaryls. I. Mechanism of the Amidation of Carboxylic Acids With Sulfuric Acid Amides," A. V. Kirsanov, Dnepropetrovsk Metallurgical Inst imeni Stalin

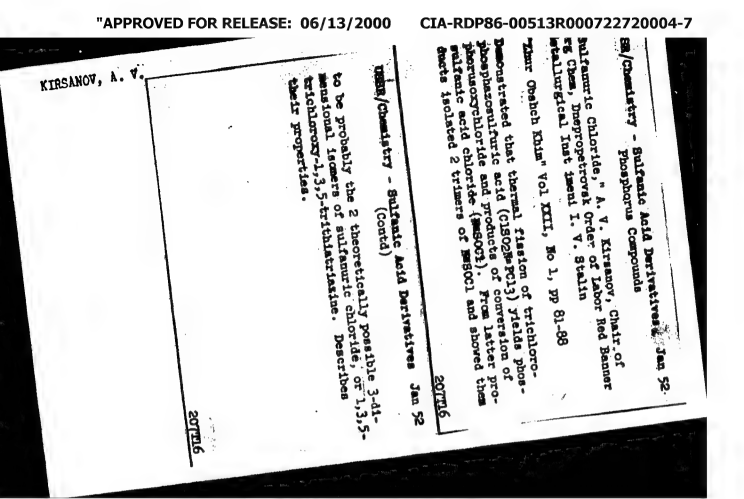
"Iz Ak Nauk SSSR, Otdel Khim Nauk" No 4, pp 710-720

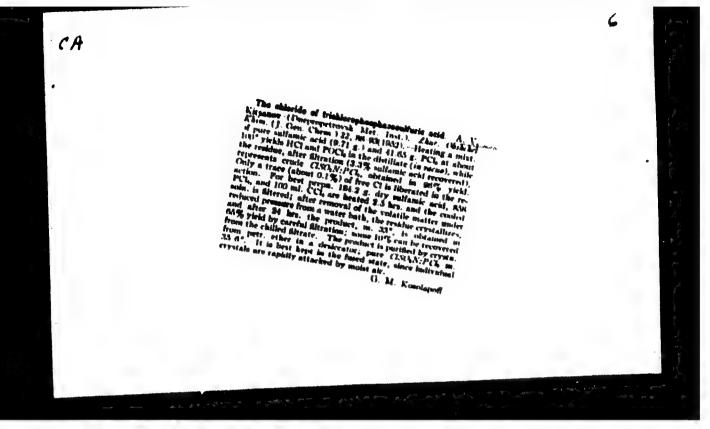
In the reaction of phenylsulfamic on carboxylic acids in pyridine, anilides of carboxylic acids are formed. The action or amines on

(1)

(CA 47 no.19:990+ 53)

acylsulfamic acids results in acylamines. Sulfrand particular acid form polypeptides of particular acid form polypeptides of people sphazosulfonaryls from aliali salts of ary sulfonic acid and the corresponding trichalogeno compds of phosphorus was found. Hydrolysis and acidolysis of trichlorophosphazosulfonaryls were studied; all theoretically possible intermediate of compds of this type were obtained. Investigated aninolysis and arylsulfonizophosphoric acids, salts of trichlorophosphazosulfonizes of arylsulfonizidophosphoric acids, chlorides of arylsulfonizidophosphoric acids, chlorides of dismilidearylsulfonizidophosphoric acids, chlorides of dismilidorarylsulfonizidophosphoric acids, and trianilide of arylsulfonizidophosphoric acids. Investigated the reactions of trichlorophosphazosulfonaryls will alcs and phenols. Obtained triesters of arylsulfonzatidophosphoric acids and diesters of arylsulfonzatidophosphoric acids and triesters of arylsulfonzatidophosphoric acids and diesters of arylsulfonzatidophosphoric acids.	KIRSANOV, A. V.		Card 5 0	r 2
	(E) (E)	arylsulfonsmidophosphoric srylsulfonimidophosphoric s reactions of trichloroph cs and phenols. Obtained nimidophosphoric acids and namidophosphoric acids.	acylsulfamic acids results in acylamines. Sulfr-44- and p-aminobenzoic acid form polypeptides of p- genophosphazosulfonaryls from alkali salts of aryl- sulfonic acid and the corresponding trichalogeno compds of phosphorus was found. Hydrolysis and acidolysis of trichlorophosphazosulfonaryls were studied; all theoretically possible intermediate and final products of hydrolysis and acidolysis of compds of this type were obtained. Investi- gated aminolysis and arylaminolysis of trichloro- phosphazosulfonaryls. Obtained triamides of aryl- sulfonimidophosphoric acids, salts of diamides of arylsulfonimidophosphoric acids, chlorides of di- anilidoarylsulfonimidophosphoric acids, diamilides (2)	





KIRSANOV, A. V.

UBSR/Chemistry - Sulf mides

Jan 52

"Hydrolysis of 2-Meth limidosulfamide and Monomethylsulfamide," A. '. Kirsanov, Yu. M. Zolotov, Chair of Org Chem, Dn propetrovsk Metallurgical Inst imeni I; V. Stal.n

"Zhur Obsheh Khim" Vol XXII, No 1. pp 151-153

Found that hydrolysis of 2-methylimidosulfamide (I) goes much more rapidly than hydrolysis of free imidosulfamide. Proposed that greater rate of hydrolysis of I is due to large distance between central N atom and neighboring 8 atoms and that stability of sulfamide ion in aq solns is due to shortening of this distance. Prepd and describes

KIRSANOV, A. V.

USSR/Chemistry - Organophosphorus Compounds

Feb 52

"Trichlorophosphasosulfonoaryls," A. V. Kirsanov, Chair of Org Chem, Dnepropetrovsk Order of Labor Red Banner Metallurgical Inst imeni I. V. Stalin

"Zhur Obshch Khim" Vol XXII, No 2, pp 269-273

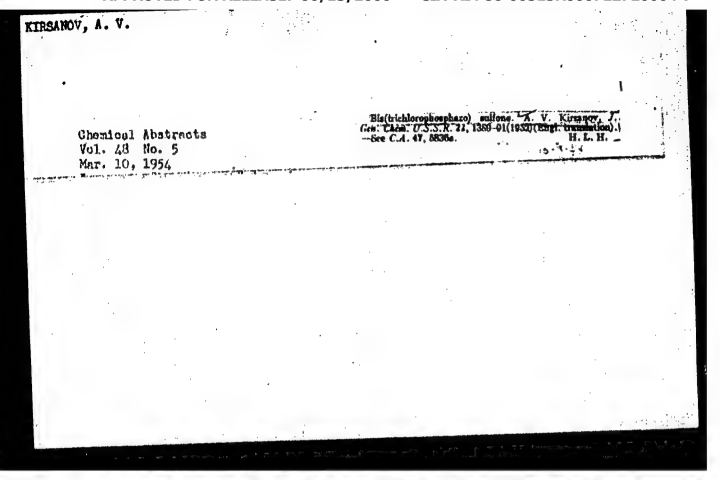
Finds that reaction of PCl₅ with arylsulfamides goes practically to completion to form trichlorophosphazosulfonoaryls (ArSO₂N=PCl₃), contrary to previous published data. Describes certain trichlorophosphazosulfonaryls.

209123

"APPROVED FOR RELEASE: 06/13/2000 CIA-RDP86-00513R000722720004-7 KIRSANOV, A. V. conversion of chlorides of carboxylic acids into clusions and verifies by expt new method for somifono-p-benzoylchloride into chloride of p-ben Discusses mech of conversion of trichlorophosphsby author. fonoaryle (ArSO2M=PCl3), which can be prepd chesply their nitriles by action of trichlorophosphasosul-USER/Chemistry - Organophosphorus somitrilosulfonic acid. Proposes on basis of con-"Zhur Obsheh Khim" Vol XXII, No 2, pp 274-278 (and conveniently) by method developed previously boxylic Acid Into the Corresponding Mitriles, W. Kirsanov, Chair of Org Chem, Dnepropetrovsk fetallurgical Inst imeni I. V. Stalin New Method for Conversion of Chlorides of Car-- American Organophosphorus Carboxylic Acid Mitriles Compounds Compounds (Contd) 7 8 X

APPROVED FOR RELEASE: 06/13/2000 CIA-RDP86-00513R000722720004-7"

700 32



KIRSANOV, A. V.

232726

USER/Chemistry - Amides

Sep 52

"Trianilide of Trimesic Acid," A. V. Kirsanov, M. L. Yegorova, Chair of Org Chem, Dnepropetrovsk Metallurgical Inst imeni I. V. Stalin

"Zhur Obshch Khim" Vol 22, No 9, pp 1614, 1615

The trianilide of trimesic acid was obtained by direct phenylamidation of trimesic acid. This product was found to melt at 320-321° and not at 118-120° as previously reported by Curtins.

232126

"APPROVED FOR RELEASE: 06/13/2000

CIA-RDP86-00513R000722720004-7

Vol. 48 No. 8 Apr. 25, 1954 Organic Chemistry

Phonylamidation of carborrile acida A. V. Kirainovi (1052) Engl. translation) — See C.A. 47, 80415.

11. L. 11.

KIRSANOV A. V.

230137

USSR/Chemistry 06913/2006 Approved FOR RELEASE: 06/13/2006 Approve

"3-Ethylimidosulfamide and Ethylsulfamide," A. V. Kirsanov and Yu. M. Zolotov, Chair of Org Chem, Dnepropetrovsk Metallurgical Inst imeni I. V. Stalin.

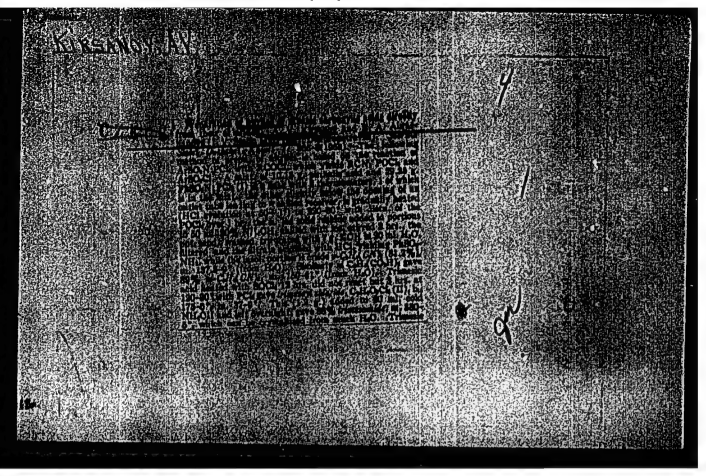
"Zhur Obshch Khim," Vol 22, No 11, pp 2038-2040

3-ethylimidosulfamide was obtained by the action of ethyl iodide on the silver salt of imidosulfamide and by the action of diazoethane on free imidosulfamide. Its properties were described. By the hydrolysis of 3-ethylimidosulfamide, ethylsulfamide was obtained, and its properties were also described.

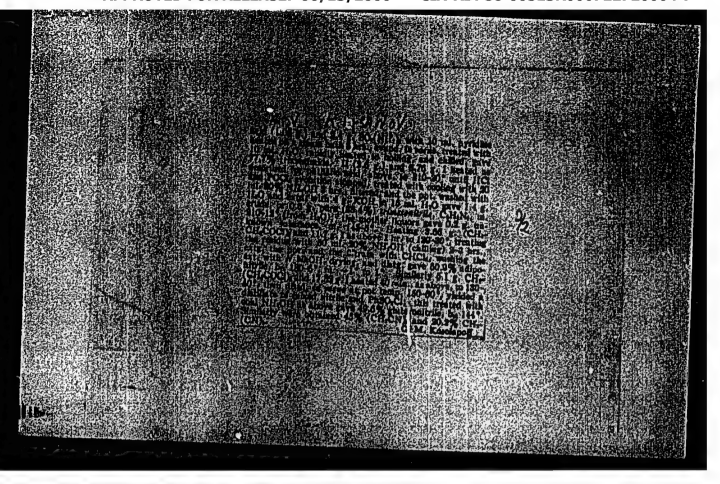
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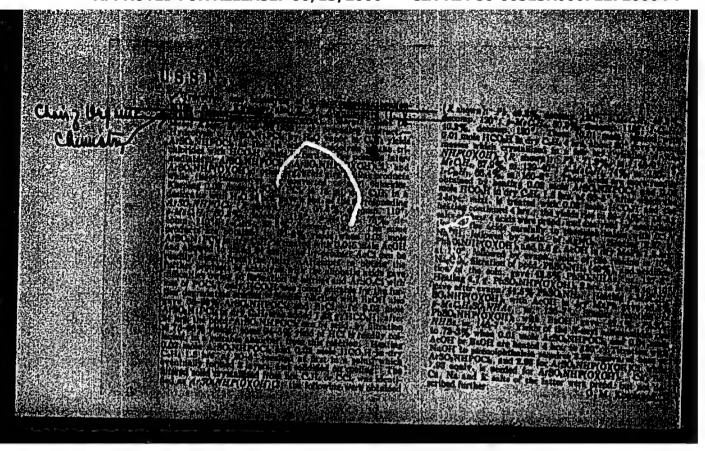
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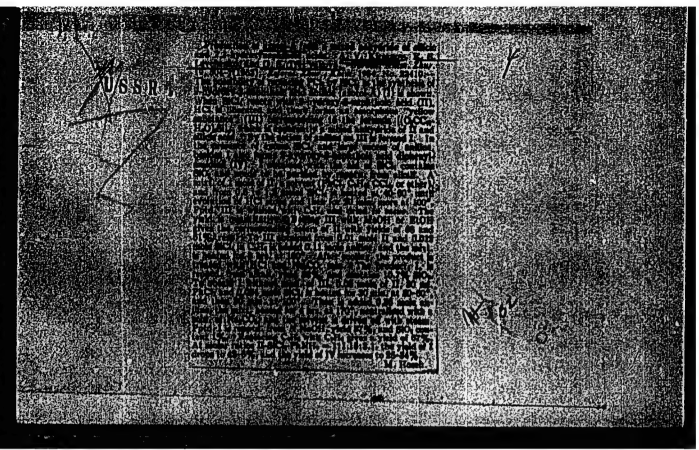
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"APPROVED FOR RELEASE: 06/13/2000 CIA-RDP86-00513R000722720004-7







KIRSANOV, A.V.; IMVCHENKO, Ye.S.; TRET YAKOVA, G.S.

Diphenylamidination of carboxylic acids. Ukr.khim.shur. 19 no.6:622-630 153. (MIRA 8:5)

1. Institut organicheskoy khimii Akademii nauk USSR (Amidines) (Acids, Fatty)

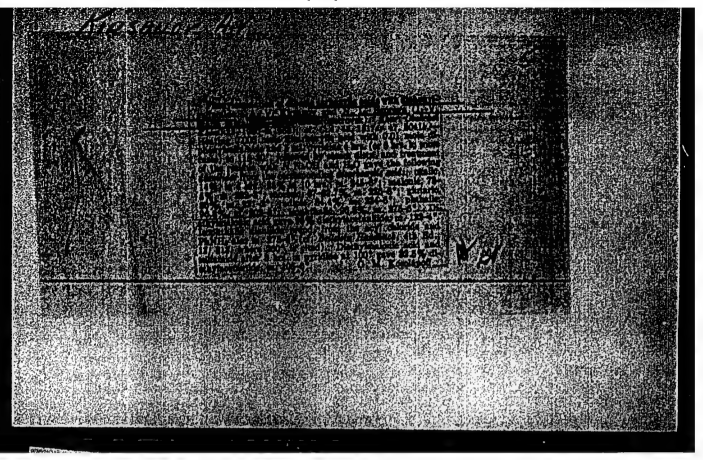
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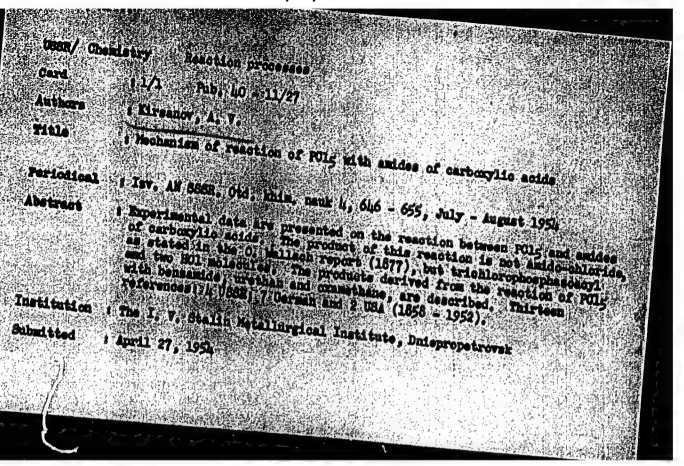
Chinical Abota
Vol 48 No 5
Mar 10, 1959
Organic channestry

MINUNIACA WILL.

Mechanism of amidating of carbarylic acide by amides of contracted. A. V. Reinov, U. V. Salpi Mer. Inst. Desproyetrows). Zast Ushakel Kasm. 23, 223-01123); C. C.A. 47, Bills.—A mechanism is proposed for the amidation of carboxylic acids with SO₂(Nils). Heating 2.75 g. MenNSO₂(NPh). 1.22 g. BrOH, and 5 ml. pyridine 4 hrs. at 516°, everyg. in suce. and treating with 15 ml. N NaOH and BtoO. gave 90.3% MenNSO₂(NPh) and 98.2% BrOH. Similar reactions with AcOH. Bit, NSO₂(NPh), and 90.4(NMe₂) gave Mentical results. Heating 2 g. MenNSO₂(NHPh), 1.22 g. BrOH, and C. ml. pyridine 3 hrs. at 115°, rvap. in suce. freating with BtoO and 2N HCL and spg. the mixed amides by even. with petr. ether gave 0.65 g. (33%) BanHPh, m. 101-2°, and 60.3% BanMer, m. 30-41°. A similar reaction with p-O₂(NC₂I₂CO₃) BrNMer, m. 30-41°. A similar reaction with p-O₂(NC₂I₃CO₃) BrNMer, m. 33-5°. Heating 18.6 g. PhNHs to 170°, adding 9.7 g. H₁(NSC₃II) or 15 min., and heating 5 min. gave 30.4% PhNHSU₃(NII). The 11.9 g.) heated 2 hrs. at 115° with 1.67 g. p-O₃(NC₃I₃CO₃) heated 2 hrs. at 116° with 1.67 g. p-O₃(NC₃I₃CO₃) heated 2 hrs. at 116° min. pyridine 6 hrs. at 115° gave 1.78 g. p-O₃(NC₃I₃CO₃) heated 2 hrs. at 116° min. pyridine 6 hrs. at 115° gave 1.78 g. p-O₃(NC₃I₃CO₃) heating 0.96 g. SO₃(NH₃), 1.86 g. PhNH₃, and 5 ml. pyridine 6 hrs. at 115° gave 1.78 g. p-O₃(NC₃I₃CO₃I). Heating 1.96 g. p-O₃(NC₃I₃CO₃I), and 5 ml. pyridine 6 hrs. at 115° gave 1.50° y. SO₃(NH₃I₃), 1.36 g. PhNH₃, m. 109-11°, and some H₃N-yields are obtained but not as high as suggested by Paqmin (C.A. 43, 2673c). Heating 1.60° g. p-O₃(NC₃I₃CO₃I), Heating 1.60° g. p-O₃(NC₃I₃CO₃I), Heating 1.60° g. p-O₃(NC₃I₃CO₃I). Heating 1.60° g. p-O₃(NC₃I₃CO₃I), Heating 1.60° g. p-O₃(NC₃I₃CO₃I), Heating 1.60° g. p-O₃(NC₃I₃CO₃I), Heating 1.60° g. p-O₃(NC₃I₃CO₃I), and heating 2 hrs. at 115° gave 1.51° gas beating 2 hrs. at 120° gave 70.4% (based on

"APPROVED FOR RELEASE: 06/13/2000 CIA-RDP86-00513R000722720004-7





TIPSANOV

USSE/Chemistry - Amidation

Card 1/1 Pub 151 20/36

Authors Kirsanov, A. V. and Abrashanova, B. A.

Title Amidation of carboxylic acids with smides of phosphoric acid

Periodical : Zhur ob, khim 21./1, 120-122, Jan 1954

The reaction of p-nitrobenzoic acid with trianilide, tri-p-toluidide and dis-nilide of phosphoric acid, and with trianilide, disnilide, trianide and mono-amide of phenylsulfonicimide phosphoric acid, was investigated. Only triani-lide, among the above mentioned phosphoric acid amides, was found to be a phenyl-amidation agent. It was also established that trianilide of phosphor-ic acid can be successfully applied for direct phenylamidation of carboxylic acids of the fatty and aromatic series. Two USSR references (1949-1953).

Institution : The I. V. Stalin, Order of Red Banner Metallurgical Institute, Faculty of Organic Chemistry, Dnepropetrovsk

Submitted : June 20, 1953

Abstract

KIRSANO

Actin Chamistry - Reaction products

Card 1/1 Pub. 151 - 21/36

Kirsanov, A. V. Jand Zolotov, Tu H. Authors

Title Tribromophes phasosulfonaryla

Periodical | Zhur, ob, khim, 24/1, 122-124, Jan 1954

Abstract The derivation of tribrosophosphasosulfonaryls from the reaction of chioramide sodium salts of sulfo solds with phosphorus tribromide, is briefly explained. The physico-chemical properties of tribromophosphasosulfonaryls, are described. The derivation and properties of tribromophosphasosulfone-phenyl, o-tolyl, p-tolyl, alpha-naphthyl and beta-naphthyl are mentioned. One USSR reference (1953).

The I. V. Stalin, Order of Red Banner Metallurgical Institute, Faculty of Institution :

Organic Cheststry, Dnepropetrovsk

Submitted June 20, 1953

USER/Chemistry as Patty soids

Oard 1/1 : Pub. 151 = 15/37

Authors | Eirsanov, A. V./ and Shevuhenko, V. I.

Chois Levere of E. 7 Langue of Long Control of London

Periodical | Zhur. obs kills 2/3 4/4-644, Mar 1954

The reaction between trichlorophosphasosulfone aryls and sodium alcoholates of hethyl; ethyl sid birtyl alcohols, was investigated. The derivation of numerous complete methyl; sthyl and butyl arylsulfonisidophosphases and their properties are described. The products of acid and alkaline hydrocured lysts of dominate arylsulfonisidophosphases are listed. A method is introduced for the direct synthesis of distryl arylsulfonisidophosphases from alcoholates of alcohols and trichlorophosphasosulfonaryls. The synthesis and properties of allohal and beta-trichlorophosphasosulfonaryls are described. Three references: 2-USSR and 1-Polish (1930-1953). Tables

Institution . The L. V. Stalin Metallurgical Institute, Despropetrovsk

Submitted : November 3, 1953

Abstract

Card 1/1

Authors | Kirsunov A. V

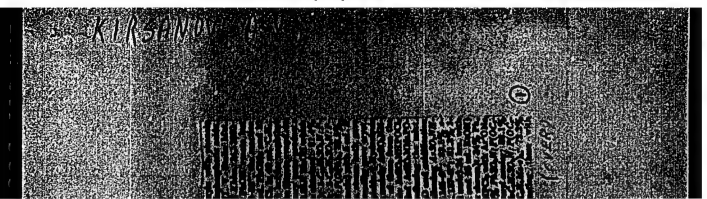
//tile | Isooyanate Phesphoryl Chloride.

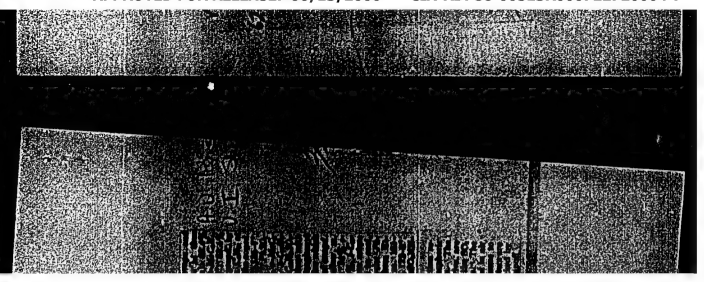
Periodical | Zhur: Ob. Khim, 24, Kd. 6, 1033 - 1038, June 1954

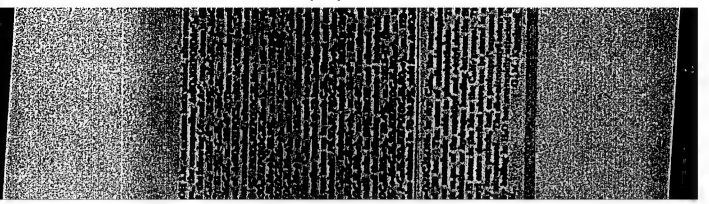
Abstract | The reaction of phosphorus pentachloride with urethan, leads to the formation of ethyl ether of trichlorophosphasocarbonic acid (athyl trichlorophosphasocarbonic acid (athyl trichlorophosphasocarbonic acid phosphoryl chloride | Ship, sther of trichlorophosphasocarbonic acid phosphoryl chloride | The physical and chemical properties of isocyanate phosphoryl chloride and ethyl ether of trichlorophosphasocarbonic acid are described | Three references; 2 German since 1852, 1877.

Institute | The I.V. Stalin-Order of the Red Banner Metallurgical Plant, Dniepropatrops.

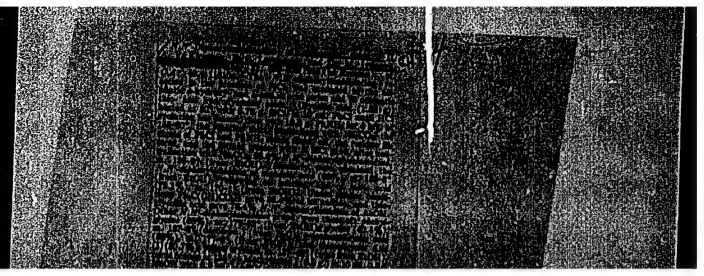
Submitted | January 3, 1954

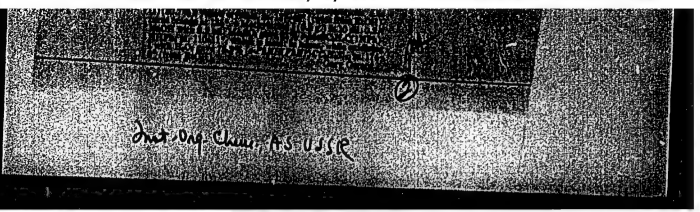


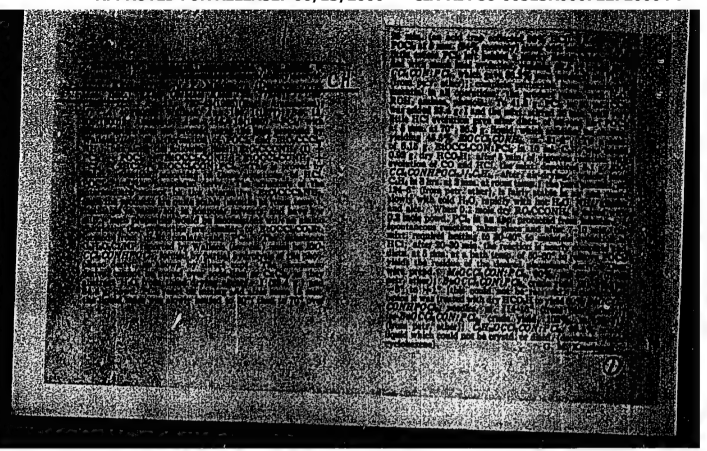












KIRSANOV, A.V.; YRGOROVA, H.L.

Dichloranhydrides of alkyl sulfonamidophosphoric acids. Zhur.ob.khim. 25 no.6:1140-1141 Je '55. (MIRA 8:12)

1. Dnepropetrovskiy metallurgicheskiy institut imeni I.V. Stalina (Phosphoric acid) (Sulfonanides)

"Reactions of Them become Pentreblers to with Acid Anales"
Taper presented at Mr. Pirst Conference on 1° sector as Consecution, in an,

Set B=3,004,841

KIRSANDVerdiver kandidat tekhnicheskikh nauk.

Selecting calculated ratios to determine the coefficient of heat convection losses in cross-tube steam flow. Sudostroenie 22 no. (MLRA 10:2)

(Boilers, Marine) (Heat-Transmission)

"APPROVED FOR RELEASE: 06/13/2000

H.V.

USSR/Organic Chemistry - Synthetic Organic Chemistry, E-2

Abst Journal: Referat Zhur - Khimiya, No 19, 1956, 61602

Author: Kirsanov, A. V., Shevchenko, V. I.

Instituteon: None

Title: Acid Chlorides of Aromatic Acid Esters of Arylsulfonimido-

Original

Periodical: Zh. obshch. khimii, 1956, 26, No 1, 250-254

There is described the reaction of the sodium salt of the chloramide of benzene sulfonic acid with acid chlorides and di-acid chlorides of the esters of phosphorous acid as a result of are formed, with good yields, the following acid chlorides (or di-acid chlorides) of aromatic esters of phenyl sulfonimidophosphoric acid (I) (listing the yield in \$): $C_{6H_5}SO_2N = PCl_2(OC_{6H_5})$ (II) (oil; 92.9); $C_{6H_5}SO_2N = PCl_2(OC_{6H_5})$ (2): $C_{6H_5}SO_2N = PCl_2(OC_{6H_5})$ (Oil; 92.9); $C_{6H_5}SO_2N = PCl_2(OC_{6H_5})$ (III) (MP 66-690; 90.1); $C_{6H_5}SO_2N = PCl_2(OC_{6H_5}CH_3)$ (V) (oil; 94.3); $C_{6H_5}SO_2N = PCl_2(OC_{6H_5}CH_3)$ (V) (oil; 96.5); $C_{6H_5}SO_2N = PCl_2(OC_{6H_5}CH_3)$ (VI)

Card 1/2

KIRSHACE HOV

USSR/Organic Chemistry - Synthetic Organic Chemistry, E-2

Abst Journal: Referat Zhur - Khimiya, No 1, 1957, 960

Author: Kirsanov, A. V., and Shevchenko, V. I.

Institution: None

Title: Reaction of Trichlorophosphazosulfonearyls with Alcohols

Original

Periodical: Zh. obshch. khimii, 1956, Vol 26, No 2, 504-510

The reaction of ArSO₂N * PCl₃ (I) with alcohols follows a number of different mechanisms and depends on the nature of the aryl radical and Abstract: on the conditions of the reaction. The first chlorine atom in I reacts faster with the alcohol than the remaining chlorine atoms; initially, an unstable molecular compound with HCl is formed which, after 60-90 minutes at 5-90, evolves 0.8 gms-equiv EC1 to form Ar802N = PC12(OR) (II); the reaction rate corresponds to the first order kinetics. When the reaction is carried out without evolution of CH1, the following mechanism is observed: II (R = CH3) + HC1 -CH3C1 + ArSO2N = : PCl2(CH). A large excess of alcohol leads to

Card 1/3

USSR/Organic Chemistry - Synthetic Organic Chemistry, E-2

Abst Journal: Referat Zhur - Khimiya, No 1, 1957, 960

Abstract: CH3ONa (prepared from 0.004 moles Na and 20 ml CH3OH) at 3-50, followed by heating for 30 minutes at 600 gives the following compounds IV (Ar and percent yield as indicated): α -CloH₇, 80.5; C₆H₅, 90.2; o-CH₃C₆H₄, 65.6. Hydrolysis of 0.01 moles II (Ar = α -CloH₇) with 100 ml water yields α-C₁₆T₇SC₂NHPO(OH)(OCH₃); the yield is 61.4%, mp 95-99°, decomposes at 1050. The compound α - $C_{10}H_7SO_2N$ = $PC1(OCH_3)_2$ (VI) is prepared by adding 0.01 moles II (Ar = α - $C_{10}H_7$) in 20 ml C_6H_6 to 0.01 mole II (Ar = α -C₁₀H₇) in 30 ml C₆H₆ and allowing the solution to stand 15 days at 26° in a dry atmosphere; the yield is 34.8%, mp 138-1400 (from benzene). Hydrolysis of 0.01 moles VI in 5 ml water at 400 with 1N NaOH, followed by acidification, yields V (Ar = α -C₁₀H₇); yield 821, mp 164-1655. The action of an excess of alcohol (10 ml) on 0.003 moles I (one hour at 200) yields from 9.6 to 37.5% IV and from 21.5 to 50.3% V. C2H5OH under the same conditions yields from 26.4 to 34.1% W and from 22.7 to 45.6% V. When the reaction is carried out over 24 hours at 200, V is formed in yields of 80%.

Card 3/3

KIRSANOV, A.V.; MERRASOVA, 2.D.

Triphonylphosphasosulfonaryls. Zhur.ob.khim. 26 me.3:903-904 Kr 156. (MLRA 9:8)

l, Ineprepetrovskiy ordena Trudevege Krasnoge snameni metallurgicheskiy institut imemi I.V. Stalina. (Sulfonaryls)

Reaction of phosphorus pentachloride with carboxylic acid anikes.

Trichlorophosphasoacyls. Zhur.ob.khim. 26 no.3:907-914 Mr '56.

(MEA 9:8)

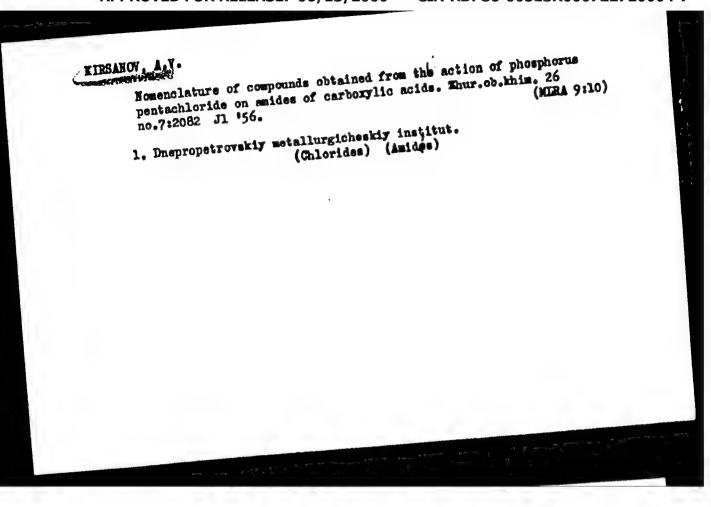
1. Institut organicheskoy khimil Akademii nauk Ukrainskoy SSR.

(Phosphorus pentachloride) (Trichlorophosphasoacyl)

Trichlorophosphasotrichloroscetyl and //-phosphoryl chlorids of trichloroiminoscetyl Enur.eb.khim. 26 no.7:2009-2014 Jl '56e trichloroiminoscetyl Enur.eb.khim. 26 no.7:2009-2014 Jl '56e (MRA 9:10)

1. Dnepropetrovskiy metallurgicheslkiy institut.

(Acetyl chloride) (Phosphorus compounds)



KIRSANOV, A.V.; LHYCHENEO, I.S.

Chlorides and esters of arylcarbamidophosphoric acids. Zhur. ob.

(NURA 10:11)

khim. 26 no.8:2285-2289 Ag 156.

1. Institut organicheskoy khimii AN USER.

(Phosphoric acid)

Acid chlorides and esters of urethanphosphoric acids. Zhur.
ob.Krim. 26 no.912642-2648 S '56.

1. Institut organicheskoy khimii Akademii nauk Ukrainskoy
SSR.

(Chlorides) (Urethamphosphoric acid)

KIRSANOV, A. V. (Dnepropetrovsk Metallurgical Inst.)

"Reaction of Pentachloride of Phosphorus with Amides of Acids" (Reaktsiya pyatikhloristogo fosfora s amidami kislot)

Chemistry and Uses of Organophosphorous Compounds (Khimiya i primeneniye fosfororganicheskikh sayedneniy), (Khimiya i primeneniye fosfororganicheskikh sayedneniy), Trudy of First Conference, 8-10 December 1955, Kazan, Trudy of First Conference, 8-10 December 1955,

485

AUTHORS:

Kirsanov, A. V., and Makitra, R. T.

TITLE:

Triaroxyphosphazosulfonearyls and Diaryl Esters of Arylsulfonamidephosphoric Acids (Triaroksifosfazosul'fonarily i diarilovya efiry arilsul fonamidofosfornykh kislot)

PERIODICAL:

Zhurnal Obshchey Khimii, 1957, Vol. 27, No. 1, pp. 21,5-248 (U.S.S.R.)

ABSTRACT:

Numerous chloro- and nitrosubstituted triaroxyphosphazosulfonaryls and diaryl esters of arylsulfonamidephosphoric acids were synthesized for the purpose of studying their insecticide properties. The o- and p-trichlorotriphenoxyphosphazosulfonaryls are described as colorless, crystalline substances (with the exception of otrichlorotriphenoxyphosphazosulfone-p-tolyl which is in liquid state at room temperature), of neutral nature, insoluble in water, easily soluble in acetone, ethyl acetate and ethyl ethers, crystallizes in ethyl alcohol, petroleum ether, and carbon tetrachloride. The o- and p-trinitrotriphenoxyphosphazosulfonaryls represent bright-yellow crystalline substances of neutral nature, well soluble in dioxane at room temperature, crystallize in benzene, insoluble in water and in a majority of organic solvents. Dinitrodiphenyl esters of arylsulfonamidophosphoric acid are

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485

Triaroxyphosphazosulfonearyls and Diaryl Esters of Arylsulfonamidephosphoric Acids

crystalline bright-yallow substances, well soluble in alcohol and acetone, almost insoluble in water and in a majority of organic solvents, crystallize in aqueous alcohol, and have the characteristics of strong monobasic acids. When mixed with alkalis, or aryls, they yield brightly-colored well-crystallizing salts.

Two tables. There are two Slavic references.

ASSOCIATION:

Academy of Sciences Ukrainian SSR, Institute of Organic Chemistry (Institut Organicheskoy Khimii, Akademii Nauk Ukrainskoy SSR)

PRESENTED BY:

SUBMITTED:

Feburuary 14, 1956

AVAILABLE:

Card 2/2

Kirsanov, A. V., and Makitra, R. G.

79-2-37/58

AUTHORS:

N-Acylemidophosphoric Acids (N-atsilamidofosfornyye Kisloty)

TITLE PERIODICAL: Zhurnal Obshchey Khimii, 1957, vol 27, No 2, pp. 450-452 (U.S.S.R.)

ABSTRACT:

The hydrolysis of didloroanhydrides of acylamidophosphoric acid yielded ten (10) acylamidophosphoric acids as listed in the table. The products are described as crystalline substances, easily soluble in water and alcohol, insoluble in acatone, ether, bensene and a majority of ether organic solvents. When heated to a melting point, all acylamidopnosphoric acids decompose and in some instances the decomposition is accompanied by indecompose and in some instances the decomposition is accompanied by in-tensive darkening and liberation of gases. They cannot be recrystallized and become useless. N-acylemidophosphoric acids are strong acids displacing carbonic and acetic acids from their salts. They submit to titration with methyl orange as monobasic and with phenolphthalein as dibasic

Card 1/2

1 table. There are 4 references of which 3 are Slavic.

"APPROVED FOR RELEASE: 06/13/2000

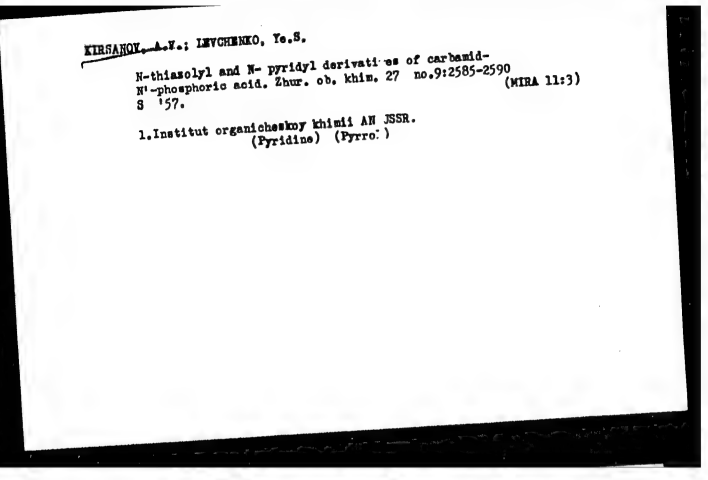
CIA-RDP86-00513R000722720004-7

ASSOCIATION: Academy of Sciences of Ukrainian SSR, Institute of Organic Chemistry

PRESENTED BY:
SUBMITTED: February 14, 1956

AVAILABLE: Library of Congress

Card 2/2



KIRSANOV, A.V.; FESHCHENKO, N.G.

Trichlorophosphazosulfonenitroaryls and the products of their hydrolysis. Zhur.ob.khim. 27 no.10:2817-2820 0 157. (MIRA 11:4)

1.Institut organicheskoy khimii Akademii nauk SSSR. (Hydrolysis) (Aromatic compounds)

79-11-35/56

AUTHORS:

Kirsanov, A. V., Molosnova, V. P.

TITLE:

The Aromatic Esters of Oxamine Acid (Effry cksaminovcy

kisloty aromaticheskogo ryada).

PERIODICAL:

Zhurnal Obshohey Khimii. 1957, Voi. 27, Hr 11, Fp. 3075-3078

ABSTRACT:

Only the phenylester synthesized by Wallaca and Liebmann and obtained in small quantities was hitherto known of the aromatic esters of oxamic acid. This phenyl ester was produced by the action of phosphorus pentachlaride upon the the athyl of examic acid with subsequent pleasi treatment on the resulting raw product. Wallach's assumption that an amidochloride as intermediate product plays an important part on that occasion has not been verified. Recent investigations say that the intermediate products are no amidochlorides, but amides of dichloralkoxyacetic acids and trichlorophosphazodichloralkoxyacetyls. Thus the formula of the formation of oxamine esters can only be represented in the following manner:

Card 1/3

The Aromatic Esters of Oxamine Acid

79-11-35/36

NH2COCCE2OR+ROH -NH2COC -OR - RCC+ ON H2COCOOR

This process of reaction has to be further investigated, as only the phenyl ester was hitherto known. In contrast to the action of aliphatic alcohols the phenois, as tests show, action of aliphatic alcohols the phenois, as tests show, act in the above-mentioned manner. By their action upon the amides of dichicralkoxyacetic acids it was possible to the amides of dichicralkoxyacetic acids it was possible to synthesize quite a number of arcmatic esters of examic acid synthesize quite a number of accompletely confirms the authors with good yields, which fact completely confirms the authors theoretical assumption. Thus it is shown that aromatic esters theoretical assumption. Thus it is shown that aromatic esters of examic acid form in the reaction of the ph. nols upon the oreaction acid form in the reaction of the ph. nols upon the amides of dichloralkoxyacetic acids and that this reaction amides of dichloralkoxyacetic acids and that this reaction amides of dichloralkoxyacetic acids and that this reaction amides of dichloralkoxyacetic acids in the ph. nols upon the foreign acids and that this reaction amides of dichloralkoxyacetic acids it was possible to acids and that this reaction amides of dichloralkoxyacetic acids it was possible to acids and that this reaction amides of dichloralkoxyacetic acids it was possible to acids and that this section acids are production of the ph. nols upon the ph. nols upon the production of the ph. nols upon the production of the ph. nols upon the ph. n

Card 2/3

The Aromatic Esters of Oxamine Acid

79-11-35/56

ASSOCIATION: Depropetrovsk Metallurgical Institute

(Dnepropetrovskiy metallurgicheskiy institut).

October 28, 1956 SUBMITTED:

Library of Congress AVAILABLE:

Oxamine acids - Derivatives 2. Aromatic esters -Production

Card 3/3

Dialkamides of Triaroxyphosphorosulphuric Acids and Aromatic Esters of the N,N-dialkylsulfamic - N' - Phosphoric Acids (Dialkilamidy triaroxsifosfazosernykh kislot i aromaticheskiye (Dialkilamidy triaroxsifosfazosernykh kislot). PERIODICAL: Zhurnal Obshchey Khimii 1957, Vol. 27, Nr 12, pp. 3241-3248 (USSR) ABSTRACT: Up to now dialkylamides of the triaroxyphosphazosulphuric acids and the esters of N,N - dialkylsulfamic - N - phosphoric acids were unknown. The present work describes their synthesis acids were unknown. The present work describes their synthesis acids were unknown. The dimethyl-and diethylamides of the and their properties. The dimethyl-and diethylamides of the action triaroxyphosphazosulphuric acids were obtained by the action of dimethylamide of the same acid on sodiumarylates according to the pattern: P2NSO2N = PCl3 + 3 NaOAr -> 3 NaCl + R2NSO2N = P(OAr) 3 These dialkylamines are colourless, crystalline, low melting and when melting non-decomposing compounds, with the exception and when melting non-decomposing compounds, with the exception of diethylamide which is liquid at room temperature. From the otherwise point of view these dialhylamides are neutral and ohemical point of view these dialhylamides are neutral and		79-12-15/43
Esters of the diskylamides of the triaroxyphosphazosulphuric (USSR) ABSTRACT: Up to now diskylamides of the triaroxyphosphazosulphuric acids and the esters of N,N - diskylsulfamic - N - phosphoric acids were unknown. The present work describes their synthesis acids were unknown. The diskylsulfamic of the and their properties. The dimethyl-and diethylamides of the and their properties. The dimethyl-and diethylamide acid and of dimethylamide of the trichlorphosphazosulfuric acid and of dimethylamide of the same acid on sodiumarylates according to the pattern: P2NSO2N = PCl3 + 3 NaOAr -> 3 NaCl + R2NSO2N = P(OAr)3 These diskylamines are colourless, crystalline, low melting and when melting non-decomposing compounds, with the exception and when melting non-decomposing compounds, with the exception of diethylamide which is liquid at room temperature. From the	AUTHORS:	Kirsanov, A. V., and Nekrasova, Z. D. Kirsanov, A. V., and Nekrasova, Z. D. Necrosic Acids
Up to now dialkylamides of the triaroxyphosphazosulphuric acids and the esters of N,N - dialkylsulfamic - N - phosphoric acids and the esters of N,N - dialkylsulfamic - N - phosphoric acids and their properties. The dimethyl-and diethylamides of the and their properties. The dimethyl-and diethylamides of the action triaroxyphosphazosulphuric acids were obtained by the action of dimethylamide of the trichlorphosphazosulfuric acid and of diethylamide of the same acid on sodiumarylates according to the pattern: P2NSO2N = PCl3 + 3 NaOAr -> 3 NaCl + R2NSO2N = P(OAr)3 These dialkylamines are colourless, crystalline, low melting these dialkylamines are colourless, with the exception and when melting non-decomposing compounds, with the exception of diethylamide which is liquid at room temperature. From the	TITLE:	(Dialkilamidy triaroksifosfazosernykh kislot).
acids and the new unknown. The present work to the acids were unknown. The dimethyl-and diethylamides of the and their properties. The dimethyl-and diethylamide by the action triaroxyphosphazosulphuric acids were obtained by the action of dimethylamide of the trichlorphosphazosulfuric acid and of dimethylamide of the same acid on sodiumarylates according the diethylamide of the same acid on sodiumarylates according to the pattern: P2NSO2N = PCl ₃ + 3 NaOAr -> 3 NaCl + R2NSO2N = P(OAr) ₃ These dialkylamines are colourless, crystalline, low melting these dialkylamines are colourless, with the exception and when melting non-decomposing compounds, with the exception of diethylamide which is liquid at room temperature. From the	PERIODICAL:	(USSE)
Card 1/3 chemical point 32	ABSTRACT:	acids and the acids were unknown. The present work and diethylamides of the acids were unknown. The dimethyl-and diethylamides of the action and their properties. The dimethyl-and diethylamide by the action triaroxyphosphazosulphuric acids were obtained by the action triaroxyphosphazosulphuric acid and of dimethylamide of the trichlorphosphazosulphuric acid and of dimethylamide of the same acid on sodiumarylates according the diethylamide of the same acid on sodiumarylates according to the pattern: P2NSO2N = PCl3 + 3 NaOAr -> 3 NaCl + R2NSO2N = P(OAr) 3 P2NSO2N = PCl3 + 3 NaOAr -> 3 NaCl + R2NSO2N = P(OAr) 3 These dialkylamines are colourless, crystalline, low melting these dialkylamines are colourless, crystalline, low melting these dialkylamines are colourless, crystalline, low melting the action and their properties.
	Card 1/3	chemical point of view

Dialkamides of Triaroxyphosphorosulphuric Acids and Aromatic Esters of the N,N-dialkylsulfamic - N' - Phosphoric Acids.

79-12-15/43

in boiling water very difficulty saponifiable products. With only few exceptions they can be saponified only by heating only few exceptions they can be saponified only by heating of alkalilyes on which occasion, however, saponification with good yields occurs only to the esters of the N,N-dialkylgood yields occurs only to the esters of the N,N-dialkylgoliamic - N- phosphoric acids (see pattern 2). They are insulfamic - N- phosphoric acids (see pattern 2) acetone, chlorosoluble in water, however, easily soluble in acetone, chlorosoluble in water, however, easily soluble in acetone, chlorosoluble in diarylform, dichlorethane, benzene and hot alcohol. The diarylform esters of the N,N-phosphoric acids were produced from the disconnection of the N,N-phosphoric acids according to the formula:

R₂NSO₂NHPOCl₂ + 2 ArONa - 2 NaCl + R₂NSO₂NHPO(OAr)₂

The diesters obtained in this way agreed with those which synthesized by saponification of the dialkylamides of the triar-oxyphosphazoacids which indicates at the presumed structure. There are 3 references, 3 of which are Slavic.

Card 2/3

Dialkamides of Triaroxyphosphorosulphuric Acids and Aromatic Esters of the N, N-dialkylsulfamic - N' - Phosphoric Acids.

79-12-15/43

ASSOCIATION:

Dnepropetrovsk Metallurgical Institute

(Dnepropetrovskiy metallurgicheskiy institut)

SUBMITTED:

October 8, 1956

AVAILABLE:

Library of Congress

Cyclic compounds - Synthesis 2. Cyclic compounds Properties 3. Dialkamides of Triaroxyphosphorosulfuric acid 4. Aromatic esters of N, N-2. Cyclic compounds -

dialkylsulfamic-N'-phosphoric acids

Card 3/3

AUTHORS:

Kirsunov, A. V., Derkach, G. I.

79-12-16/43

Trichlorphosphazoaciles, Trichloroisophosphazoaciles

TITLE:

(Trikhlorfosfazoatsily, trikhlorizofosfazoatsily i ikh and Their Derivatives

proizvodnyye).

PERIODICAL:

Zhurnal Obshchey Khimii 1957, Vol. 27, Nr 12, pp. 3248-3254

ABSTRACT:

As it has been shown before trichlorophosphazoaciles of the RCON = PCl3 type are obtained by the action of pentachloride on the amides of the carboxylic acids, which on the occasion of partial hydrolisis form dichloroanhydrides of the acilamidophosphoric acids. Up to now only a trichloroisophosphazoacile was known it e. trichloroisophosphazotrichloracetyl and some corresponding triaroxyisophosphazoaciles. It is of interest whother also from other carboxylic acids trichloroisophosphazoaciles and triaroxyisophosphazoaciles occur or whether the trichloroisophosphazoaciles occur only in the case of trichloroacetic acid and its analogs. Trichlorophosphazoaciles were produced for the diphenylchloracetic acid, triphenylacetic acid and p - nitrobenzoic acid and their thermal stability was investigated. It was demonstrated that

Card 1/2

Trichlorphosphazoaciles, Trichloroisophosphazoaciles and Their Derivatives.

79-12-16/43

trichloroisophosphazoaciles occur not only for trichloracetic acid and its analogs but also for diphenylchloracetic acid and p - nitrobenzoic acid. Trichloroisophosphazotriphenylacetyl could not be obtained, since dichloroanhydride of the triphenylacetylamidophosphoric acid when thermated has no durability. The authors obtained the di- and tri - x - naphtoxyderivatives from the corresponding synthes_zed trichlorophosphazo- and trichloroisophosphazo compounds. The synthesized Tri -d- naphtoxyisophosphazo - p - nitrobenzoyl is the first derivative of trichloroisophosphazoaciles obtained from crystals.

There are 6 references, 4 of which are Slavic.

Dnepropetrovsk Metallurgical Institute ASSOCIATION:

(Dnepropetrovskiy metallurgicheskiy institut).

October 16, 1956 SUBMITTED:

Library of Congress Trichlorophosphazoaciles - Chemical analysis AVAILABLE:

Trichloroisophosphazoaciles - Chemical analysis Card 2/2

KIRSANOV. A. V.

79.1-7/63

Molosnova, V. P. Kirsanov, A. V. AUTHORS:

The Reaction of Phosphorus Pentachleride With Esters of Oxamic Acid (Reaktsiya pyatikhioristogo fosfora s efirani oksaminovoy kisloty) The Amides of Alkoxydichloroacotic TITLE Acida (Amidy alkoksidikhloruksusnykh kislot)

Zhurnal Obshchey Khimii, 1958, Vol. 28; Er 1 pp.30-35(USSR) PERIODICAL:

It was already earlier shown that under the influence of phosphorus pentachloride upon the ethyl ester of examic acid the amide of ethoxydichloroucetic acid and emichlorophespha ABSTRACT: zoethoxydichloroscetyl are consecutively obtained according

to the following scheme:

The synthesis of trichlorophosphaspharoethacharolatelland The perojection was protectly fundated and the are apean that reaction (2) is generally valid for the synthesis of

Cari 1/3

The Reaction of Phosphorus Pentachleride With Esters of Oxemia Acid . The Amides of Alkoxydichlorophosphazosikaxydichlorosasetyle. It was of interest to find out whether reaction () night class to generall, va lid and whether it should also be considered a general nother for the production of unales of alkexydichleromeans acads. It was found that reaction (1) takes place sufficiently fust at considerably lower temperatures than reaction (2) for the mothyl-, ethyl-, butyl and isobatyl-ester of examic and. This fact permitted to obtain the amides of the acrrespending alkoxydichloroacetic acids with good yields, although reaction (2) in all cases takes place besile reaction (1). Quantitatively the amides are hard to be separated from the acetyls. The yieldof amides is quite different according to the ester of examic acid used. Reaction (a) takes place near difficult. ly for the methylester, more susily for the ethylm and quite easily for the butyl- and isobutyl ester of oxamic acid. The structure of the amides of alkoxydichloroacetic acids can quite incontestably be proved by their reaction with phentis (see formulae on page 33). There are 6 references, 5 of which are Slavic.

Card 2/3

79-1-7/63

The Reaction of Phosphorus Pentachloride With Esters of Oxamic Acid. The Amides of Alkoxydichloroacetic Acids

ASSOCIATION: Despropetrovak Metallurgical Institute

(Dnepropetrovskiy metallurgicheskiy institut)

SUBMITTED:

November 20, 1956

AVAILABLE:

Library of Congress

1. Chlorides 2. Oximidines 3. Chemistry Card 3/3

CIA-RDP86-00513R000722720004-7" APPROVED FOR RELEASE: 06/13/2000

KIRSANOV. A.V.

AUTHORS: Mirsanov, A. V., Makitra, R. C. 79-1-8/63

TITLE: Diesters of Aromatic Acylamidop Coughoric Acids

(Diefiry aromaticheskikh atsilamidofo formykh kielot)

PERIODICAL: Thurmal Obshchey Khimii, 1950, Vol. 28, Nr 1, p. 3,-40

(Haur)

ABSTRACT: Like the diesters of arylaulfour idephospheric acids the

diesters of acylamidophosphoric acids should also be representable by the influence of arylates or alcoholates of sodius upon dichloraphydrides of acylamidophosphoric acids according to scheme (1), or by saponification of

triaroxyphoaphanescyls according to scheme (2) $Arconnection + 200R \rightarrow 2 Macl + Arconnection (OR)_2$ (1) $Arcon-P(OR)_2 + H_2O \rightarrow RO' + Arconnection (OR)_2$ (2)

The dimethylester of bemosylamodiphosymoric acid is obtained according to scheme (1) in the presence of much sodium methylate and methyl alcohol with a yield of 37.0%, which

is not the case with the promatic directors (10 - 265).

Carl 1,3 According to scheme (2) the diarglesters are obtained with

79-1-8/63

Diesters of Aromatic Acylamidophosphoric Asils

a good yield, where one can start directly from triculorophosphazoncyls. With dry solium arylates in a tenzene-, other- or dioxane-solution where acyle yiel; the corresponding triaroxyphogole meacyle according to the following scheme: Arcor = Poly+ 3MaON -> 3MaCl = Arcor=P(OR);

These acyls are much more easily said ified to the diesters of triaroxyphosphezoculfon ryle, so that a heating of 5 - lo minutes with mater is suffici at for attaining the complete conversion of triaroxyphosphano cyls to the diarylesters of acylamidophosphoric acids. In the synthesis of the diecters of p-ohlorobenzoylamidophosphoric acid p-chlorobenznitrile was in some cases liberated as byproduct. According to (2) the authors obtained the methyl-, phenyl-, p-cresylic, p-chlorophenyl- and p-nitrophenyl--enters of benzoyl-, p-chlorobenzoyl- and p-nitrobenzoylamidophosphoric acids as well as the di-a-naphtylester of benzoylamidophesphoric acid.

ASSOCIATION:

Laboratory for Insecticides of the Institute for Organic Chemistry A.: Ukrainian SSR (Laboratoriya insolititsidov Instituta organicheskoy khinii Akad mii nauk Ukrainsko SSR)

Card 2/3

"APPROVED FOR RELEASE: 06/13/2000 CIA-RDP86-00513R000722720004-7

Diesters of Aronatic Acylamidophosphoric Acids

79-1-8/63

SUBMITTED:

November 26, 1956

AVAILABLE: Library of Congress

Card 3/3

1. Chemistry 2. Arylates 3. Alcoholates

KIRSHNOU. AV.

"C-2-15/34

AU HOLS:

Kirsanov, A. V., Feshchenko, M. G.

TTTLE:

Trinethoxy- and Triaroxyphosphasosulfonnitrophengla and Diesters of Mitrorhenylsulfonamidophosphoric Acids (Trimetoksi- i triaroksifosfazogul'fonnitrofenily i diefiry nitrofenilsul'fon midofosfornyth

kislot)

PERIODICAL:

Zhurnal Obshehey Khimii, 1958, Vol. 23, Nr 2, pl. 359 - 343 (USSR)

ABSTRACT:

Trimethoxy- and triaroxyphosphasosulfonnitrophenyls (I) and diesters of nitrophenylsulfonamidophosphoric acids (II) were synthesized for the purpose of investigating their insecticidic properties and as a starting point for the production of N-phospholic acid derivatives of sulfanilamides. (I) was obtained by interaction of trichlorophosphasosulfonnitrophonyls (reference 1) with methylate and sudium arylates in a bennenc solution (reference 2), Trincthoxy-, triphenoxy-, tri-p-chlorotriphenoxy-, 'ri-c- and tri-p-trinitrotriphenoxyphosphasoculfone-o-, in- and p-nitrophenyls (I) (table 1) were obtained in this manner. (I) represent colorless crystalline substances of a neutral character. They do not discolve in water. Trimethoxyphosphasosulfonnitrophenyls (III) within one heur saponify on boiling with 96% alcohol to the corresponding dicaters (XIV).

Card 1/3

79-2-13/64

Trimethoxy- and Triaroxyphosphasosulfonnitrophenyls and Diesters of Mitrophenylsulfonamidophospheric Acids

Triphenoxyphosphasosulfonnitrophenyls (IV) do not change under the same conditions. Trimethoxy- and triphenoxyphosphasosulfonnitgophenyls welt at comparatively low temperatures (from 56 - 104°C). (I) do not saponify upon the influence of aqueous alkaline solutions, which is explained by their insolubility in wher. In acceous alkaline spirit solutions they easily saponify to (II). But (II) can more conveniently be produced from dichloroanhydrides of nitrophenylsulfonamidophosphoric acids (reference 1) by means of the influence of sodium arylates and - methylate in a diomene solution. Thus dimethyl-, di-p-chlorodiphenyl-di-o- and di-p-nitrodiphenyl ethers of o-, a- and p-nitrophenylaulfonamidophosphoric acids (II) (table 2) were produced. (II) can be eliminated in the form of salts, but it is more convenient in the form of free diesters. (II) represent comparatively high-melting (from 134 - 194°C), crystalline, colorless substances incoluble in water. They nelt at far higher temperatures than the corresponding (I), with the exception of two p-nitrophenolethers (XI and XIII) which aclt under the corresponding (I). (II) are monobasic acids which exactly tetrate in the presence of phenolphthalein and yield well-crystallizing and water-soluble sodium salts. It is interesting that the sodium salts of the p-chlorophenylethers of nitrophenylsulfonamidophosphoric acids on withdrawal of the water solutions by the other com-

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79-2-13/64

Trimethoxy- and Triaroxyphosphasosulfonnitrophenyls and Diesters of Hitrophenylsulfonamidophosphoric Acids

pletely go over into the ether layer. Sodium salts of other diesters are not withdrawn from the water solution by the ether. There are 2 tables, and 1 Slavic reference.

Institute for Organic Chemistry AS Ukrainian SSR ASSOCIATION:

(Institut organicheskoy khimii Akademii nauk USSR)

December 24, 1956 SUBMITTED:

Library of Congress AVAILABLE:

Card 3/3

CIA-RDP86-00513R000722720004-7" APPROVED FOR RELEASE: 06/13/2000

HIRSPNOU AV.

79-2-14/54

AUTHORS:

Kirsanov, A. V., Zolotov, Yu. M.

TITLE:

Substituted Imido- and Monoarylsulfamides (Zameshchennyye imido- i nonoarilsulfamidy)

PERIODICAL:

Zhurnal Obshchey Khimii, 1958, Vol. 28, Nr 2, pp. 343 - 347 (USSR)

ABSTRACT:

Inidosulfamide formerly was a practically unattainable substance (reference 1) and the substituted imidesulfacides were unknown. At present the imidosulfamide is absolutely attainable (reference 2) and 3-methyl- and 3-ethylimidosulfamides (reference 3) were produced from it. 1- and 1,5-substituted inidosulfamides were hitherto unknown. According to its chemical properties the free imidosulfmaide is a monobasic acid which in strength is almost not inferior to sulfuric or hydrochloric acid. The method (reference 2) proved to be suitable for the production of 1,5- disubstituted and 1,1,5,5--tetrasubstituted imidesulfamides. Thus this nethod is Weneral method for the production of as well nonsubstituted as of 1,5-diand 1,1,5,5-tetrasubstituted inidosulfamides according to the scheme $2R_2NSO_2NH_2+OH^- \longrightarrow NH_3+H_2O + (R_2NSO_2NSO_2NR_2)^-$, where R may be hydrogen, alkyls or aryls. In the case of 1,5-diphenylimidosulfamide the initially forming unstable phenyleulfaminic acid (reference 4) is at once hydrolized and forms an acid aviline-sulfate.

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75-2-14/64

Substituted Inido- and Monoarylsulfamides

The scheme suggested for the formation of the imidosulfamide by means of an alkali liquor (reference 2) is designated as false, which is proved by the formation of (I) and (II) from N,N-dimethylsulfacide and Hell-diphenyleulfamide. The production of imidesulfamide from sulfamide and tertiary bases may take place according to $\text{MII}_2\text{SO}_2\text{MII}_2 + \text{Py} \iff \text{PyH}^+ + \text{MII}_2\text{SO}_2\text{MH}^$ etc. When all the scheme schemes determined by the authors should be correct, it must be possible to obtain substituted imidosulfamides from all substituted sulfamides with the exception of tetrasubstituted ones. This conclusion is at present examined. For the synthesis of other 1,5-disubstituted imidosulfamiles it was necessary to obtain monosubstituted sulfamides of the type RMSO_NH2. These compounds were obtained by Denivell, Battegay and Meybek (reference 5) in quite a complicated manner. Paken (reference 7) produced butyl-, cyclohexyl- and piperidylsulfamides in a simpler manner by the influence of sulfnaide upon the corresponding anines. Paken does not say anything on the Production of H-monoarylaulfamides according to this scheme. Therefore the authors decided to determine the possibility of the production of N-arylaulfamides by the influence of aroundic anines

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upon sulfamides. It became evident that the aromatic amines on heating easily react with sulfamide. The yield of monoarylsulfamides

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Substituted Imido- and Monoarylsulfamides

and diarylsulfamides, however, is not large. This is easy to understand, as on the influence of amines upon sulfamide the imidosulf-amide must also be produced beside the formation of substituted sulfamides. This imidosulfamide practically represents the only reaction product in the inveraction of the sulfamide with teriary amines (reference 2). Besides the already formed M-arylsulfamide on heating andunder the influence of an amine excess may yield the corresponding 1,5-diarylimidosulfamide. At present it is impossible to find out whether the M-arylsulfamides are formed directly from the amines and the sulfamide according to the summary scheme (IV) or only as products of the hydrolysis of the 1,5-diarylsulfamide being in the reaction mixture, according to scheme (III). There are 1 table, and 7 references, 3 of which are Slavic.

ASSOCIATION:

Institute for Metallurgy, Dnepropetrovsk

(Dnepropatrovskiy metal lurgichaskiy institut)

SUBMITTED:

January 17, 1957

AVAILABLE:

Library of Congress

Card 3/3

ardanov,		79-2-15/54
UTHORS:	Kirsanov, A. V., Molosnova, V. P. Trichlorophosphasoaroxydichloracetyls (wrikhlorfosfazoaroksidikhlor-
PITLE:	Trichlorophosphasoaroxydichloracety15 (atsetily)	*** *** (USSR)
PERIODICAL:	Zhurnal Obshchey Khimii, 1950, Vol. 28,	, Nr 2, pr. 216 - 750 (obtained)
ABSTRACT:	It was recently discovered that the all react with pentachlorophosphorus and a oxydichloroacetic acids (reference 1) subsequently trichlorophosphascalkoxyd subsequently trichlorophosphascalkoxyd	ichloracetyls (reference 2)
	ROCCOCOMI + PC15 POC13 + ROCC12	CONT PC1 (II)
	For aromatic esters of Oxidate procession with not possible to limit the reaction with sence of a large excess of oxanic-acid sence of a large excess of oxanic-acid take place or it at once takes place the formation of trichlorophosphasoar aroccocons, + 2PCl ₂ -> 2HCl + POCl ₃ +	th scheme (1). Even in the scheme (1) with according to scheme (III) with according to scheme (III) with a scheme (III) with a scheme (III) with a scheme (III) and a scheme (III) and a scheme in reference 1.
	Arococomic + 2PCl ₅ \longrightarrow 2HCl + POCl ₃ + A possible explanation of this phenomerate favorable conditions the reacti	in releasence

79-2-15/64

Trichlorophosphasouroxydichloracetyls

scheme (III) quantitatively or almost quantitatively with the formation of colorless, crystalline trichlorophosphasoaroxydichloracetyls (IV). The "raw products" as a rule melt at temperatures only a few degrees lower than the recrystallized ones and are usually colorless. Only in the case of narhthoxy derivatives (XI and XII) the raw products are colored and melt at considerably lower temperatures than the pure substances. According to its physical and chemical properties (IV) is similar to tricklorophosphasoalkoxydichloracetyl (reference 2). Under the influence of water vapors or, still better, of dehydrated formic acid (IV) yield dichloranhydrides of aroxydichloracetylamidophosphoric acids (V) with a good yield according to the scheme: Arocci₂con = Pci₃ + HCOOH -> co + HCl + Arocci₂conHPoCi₂. But on heating (IV), in contrast to trichlorophosphasoalkoxyacetyls, decompose considerably more difficult. On that occasion they do not split off any chlorobenzene and considerably more difficult and slowly (only at 160°C) phosphorus chloroxide. (Y) represent colorless, crystalline substances. They melt at omederably higher tenperatures than the corresponding trichlorophosphasoaroxydichloracetyls (IV) and in the usual organic solvents they are more difficult to solve than (IV). At room temperatures (V) slowly react with

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79-2-15/64

Trichlorophosphasoaroxydichloracetyls

water. Therefore thy can be stored without any special precautionary measures. On heating with water (V) rapidly hydrolylates and energetically reacts with substances whose molecules contain active hydrogen atoms. All solvents and initial substances were most exactly dehydrated. All tests were performed in a manner that the reaction mixtures and reaction products as far as possible did not get in touch with atmospheric moisture. There are 3 references, all of which are Slavic.

ASSOCIATION:

Institute for Metallurgy, Dne propetrovsk (Dne propetrovskiy metallurgicheskiy institut)

SUBMITTED:

January 17, 1957

AVAILABLE:

Library of Congress

Card 3/3

C	Acids	(Efiry	

TITLE:

AUTHORS:

Kirsanov, A. V., Feshchenko, N. G.

Ester of Aminophenylsulfonamido-Phosphoric Acids (E

aminofenilsul'fonamidofosfornykh kislot)

PERIODICAL: Zhurnal Obshchey Khimii, 1958, Vol. 28, Hr 4, pp. 1049-1052(USSR)

ABSTRACT

Dimethyl- and diphenyl ester of c-. m., and p-aminophenylsulfonamido-phosphoric acids (formula I) were produced according to the following reaction scheme:

 $\text{HO}_2\text{C}_6\text{H}_4\text{SO}_2\text{NH}_2\xrightarrow{\text{PCl}_5} \xrightarrow{\text{NO}_2\text{C}_6\text{H}_4\text{SO}_2\text{N-PCl}_3} \xrightarrow{\text{HCOOH}} \text{HO}_2\text{C}_6\text{H}_4\text{SO}_2\text{FHPOCl}_2\xrightarrow{\text{PCl}_5}$

 $\xrightarrow{\text{2NaOR}} \text{NO}_2 \text{C}_6 \text{H}_4 \text{SO}_2 \text{NHPO(OR)}_2 \xrightarrow{\text{6 H}} \text{NH}_2 \text{C}_6 \text{H}_4 \text{SO}_2 \text{NHPO(OR)}_2$

The reduction of diesters of the nitrophenylsulfonamido-phosphoric acids was performed with hydrogen at presence of a palladium catalyst in alcoholic solution at room temperature and at a pressure of ca. 100 torr. The yields were quantitatively. The diester produced by this way form colorless crystals,

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Ester of Aminophenylsulfonamido-Phosphoric Acids

which have only weakly basic, but strong acid properties. They solve readily in aqueous soda solution and can be titrated as monobasic acids. Their aqueous solutions react acid with Congo red. The dimethyl enter of the aminophenylsulfonamido phosphoric acids are readily soluble in hot water and can be diazotized on the usual conditions (as aniline). The corresponding diphenyl ester are in hot water difficultly to solve, in alcohol more readily soluble. They can be diazotized on the same conditions as aromatic amines with electronegative substituents (e.g. nitroaniline). The amides of the comme and praminobenzenesulfonic acids, to the derivatives of which belong the compounds of the formula I, have little different melting points (153, 142, 163°). The lowest melting point has the m-isomeric, the highest the p-isomeric (Ref 3). At the corresponding isomeric compounds of the formula I on the contrary the melting points are far from each other. The highest melting point here has the m-isomeric, the lowest the consoneric. The ortho aminocompounds of the formula I are at room temperature very readily soluble in acetone, the para-isomeric only with difficulty and in case of heating. The meta-isomerics are practically

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Ester of Aminophenylsulfonamido-Phosphoric Acids

insoluble in acetone. For the strong change of the melting points as well as for the great differences of the solubility in acetone the following explanation is probable: In the case of the o- and p-isomerics intra- and intermolecular hydrogen bindings form, the formation of which in case of the m-isomerics is complicated or impossible. The influence of the hydrogen bridges upon the melting points of the amides of the aminobenzenesulfonic acids is because of absence of the polarizing influence of the phosphoric acid rest essentially lower; therefore the melting points of the isomerics are close together. For the characteristic of the compounds with the formula I their benzoyl derivatives were produced according to the method by Schotten-Baumann. These compounds form colorless crystals, which have strong acid and no basic properties. Their aqueous solutions react acid with Congo red. The melting points partly are higher and partly are lower than the melting points of the corresponding compounds of the formula I.

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In an experimental part the synthesis and the properties

Ester of Aminophenylsulfonamido-Phosphoric Acids

of the compounds of the formula I and of their benzoyl derivatives are described exactly. There are 2 tables and 4 references, 4 of which are Soviet.

references, 4 of which are poviet.

ASSOCIATION: Institut organicheskoy khimii Akademii nauk Ukrainskoy SSR

(Institute for Organic Chemistry, AS Ukrainian SSR)

SUBMITTED: February 14, 1957

Card 4/4

AUTHORS:

Kirsanov, A. V., Yegorova, N. L.

79-28-4-44/60

TITLE:

Triaroxyphosphorazosulfonalalkyls and Aromatic Esters of Alkylsulfonamidophosphoric Acids (Triaroksifosfazosul'fonalkily i aromaticheskiye efiry alkilsulifonamidofosfornykh kislot)

PERIODICAL:

Zhurnal Obshchey Khimii, 1958, Vol. 28, Nr 4, pp. 1052-1055

(USSR)

ABSTRACT:

Triaroxyphosphorazosulfonalkyls were producet by reaction of sodium arylates with trichlorophosphorazosulfonalkyls (ref 1)

in benzene solution:

 $RSO_2N=PCl_3 + 3 Arona ---> RSO_2N = P (OAr)_3 + 3 NaCl$

Sodium phenolate and sodium-p-chlorphenolate react very with trichlorophosphorazosulfonalkyls. With dry sodium phenolate without solvent, the reaction very violently takes place and the reaction mixture carbonizes. Sodium-p-nitrophenolate reacts less violently; the reaction takes place on heating in the

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water bath in the course of 2-3 hours. Yields, melting points

Triaroxyphosphorazosulfonalkyls and Aromatic Esters of Alkylsulfonamidophosphoric Acids

79-28-4-44/60

and analytical data of the compounds of formula I obtained in this way are exactly mentioned. The compounds are colorless crystallized substances (with the one exception of the liquid derivative $Ar = C_6H_5$, $R = n-C_4H_9$) which are easily soluble in acetone, dioxans, a little more difficultly in benzene, ether and alcohol. They dissolve in boiling tetrachloride and petroleum ether as well, whilst in hot water very difficultly, in cold water they are insoluble. According to their physical and chemical properties the compounds of the formula I are very similar to triaroxyphosphorazosulfcnaryls (ref 2), however, they differ by an higher solubility in boiling water and polar solvents. The compounds of formula I are neutral substances which are relatively constant against water and atmospheric moisture. An exception is represented by the derivative with Ar = p-NO₂C₆H₄, R = C₂H₅ (II), which is very easily hydrolyzable. Already in its solutions in 96 % ethanol a complete saponification takes place, where p-dinitrophenyl

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Triaroxyphosphorazosulfonalkyls and Aromatic Esters of Alkylsulfonamidophosphoric Acids

79-28-4-44/60

ester of ethylsulfonamidophosphoric acid (III) and nitrophenol are formed:

 $c_2H_5So_2N=P(oc_6H_4No_2-p)_5 + H_2O ---> c_2H_5So_2NHPO(oc_6H_4No_2-p)_2+$ + $p-No_2c_6H_4oH_2$

All the other compounds of formula I are not modified on heating with alcohol, however, under the action of caustic alkalies in alcoholic-aqueous solution they are easily saponified. In this case salts of the corresponding diaryl ester of alkylsulfonamidophosphoric acids (IV) form, the yields, melting points and analytical data of which are mentioned. They easily dissolve in acetone and hot alcohol, very difficultly in cold - more easily in boiling water. In most of the unpolar solvents they are difficultly soluble. In chemical respects they are strong monobasic acids. With respect to hydrolysis in an acid, as well as in an alkaline

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Triaroxyphosphorazosulfonalalkyls and Aromatic Esters of Alkylsulfonamidophosphoric Acids

79-28-4-44/60

medium they are essentially more constant than the compounds of formula I. There structure not only unequivocally results from the formation by saponification of the compounds of formula I_{ν} but was also proved by the synthesis of diacetic chlorides of alkylsulfonamidophosphoric acids with sodium arylates:

RSO₂NHPOCl₂ + 2Arona -> RSO₂NHPO(OAr)₂+ 2NaCl

Syntheses and analytical data of the mentioned compounds are exactly described in an experimental part.

There are 2 tables and 3 references, 3 of which are Soviet

ASSOCIATION:

Dnepropetrovskiy metallurgicheskiy institut (Dnepropetrovsk Metallurgical Institute)

SUBMITTED:

February 4, 1957

Card 4/4

AUTHORS:

Kirsanov. A. V. Derkach, G. I.,

79-28-5-21/69

Makitra, R. G.

TITLE:

Triaroxyphosphazoacyl (Triaroksifosfazoatsily)

PERIODICAL:

Zhurnal Obshchey Khimii, 1958, Vol. 28, Nr 5,

pp. 1227-1232 (USSR)

ABSTRACT:

The similarity of truaroxyphosphazoacyls (I) and truaroxyphosphazosulfone-compounds (II) shows up in a number of common chemical properties so that their reactions of for-

mations are in common: $RSO_2N = PC1_3 + 3NaOR! \longrightarrow 3NaC1 + RSO_2N = P(OR!)_3$

 $RCON = PC1_3 + 3NaOR! \longrightarrow 3NaC1 + RCON = P(OR!)_3$

Between them, however, also specific chemical differences with regard to heating and hydrolysis. The compounds (I) split off rather easily (depending on the radical) from the corresponding triesters of phosphoric acid and produce nitriles according to the scheme RCON = P(OR') OP(OR')

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+RCN (III). The compounds (II) are very much stable against

Triaroxyphosphazoacyl

79-28-5-21/69

heating so that until now there has been no case of splitting according to scheme (III). They saponify easily with alkali liquors under the formation of salts of the diesters of the corresponding alkyl- or aryl--sulfonamidophosphoric acids, but they do not saponify with water in neutral solutions. Therefore the synthesis and the separation of the products (II) do not meet with any difficulties because of the easy saponifiability. All compounds (I) saponify on boiling practically quantitatively to the diesters of the acylamidophosphoric acids for which reason the synthesis, separation and purification of the triaroxyphosphazoacyls takes place so difficulty; for the same reason in the experiments care must be taken that they do not come into contact with atmospheric humidity. This difference can apparently be explained by the fact that in the saponification of the compounds (I) in alkali solutions the carbon- and oxygen atoms of the carboxyl-group take part in the hydrolysis and increase the positive charge of the phosphorus atom according to the given scheme 1. In the saponification of the compounds (II) mainly only the n.trogen- and phospho-

Card 2/3

Triaroxyphosphazoacyl

79-28-5-21/69

rus atoms take part in the hydrolysis (see scheme 2). There are 2 tables and 7 references, 6 of which are

Soviet.

ASSOCIATION: Institut organicheskoy khimii AN Ukrainskoy SSR

(Institute for Organic Chemistry, AS Ukrainian SSR)

SUBMITTED: February 22, 1957

Card 3/3

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. (b. 1. 1.)	11 change at 10 0000 , take 301/1/- 3-5-5/65	
. wiles	Anilitin of allighbullon milophosphoric acids (aniliay allike alfon miloposfora, kh kislot)	
1 .::13013L:	Zhurn1 obshahe/ khimii, 1950, Vol. 20, br 6, pr. 1507-153/ (USSA)	
192 UT :	On the action of animine on trichlorophos, he zesulfondkyls the formation of animidedichlorophosphanosulfondlight of the type $30_2a = 101_2$ (MiG $_6H_5$) (1), of disnilided derophos-	
	phazosulfonalkyls of the type02 N=FC1(IHC/H5)2	
	and of wrightline hough zonal fondkyla of the life and as all could as a profess (1) could	
	not be obtained (Ref. 2). The compounds (il) are obtained in sufficiently good yields in the conversion of trichloro hosphacosulfonalkyls with aniline in carron tetrachloride. They are crystalline substances of heated character and hydrolize easily to disnilides of the alkylsulfon vidophosphoric acids (III) on heating their	
Jack 1/3	solutions in 955 alcohol or in boilin water according to	

anilides of Alkylaulfonemidophospa@ric Acids

301/13-28-6-14/63

the scheme $RSO_2H = FCI(RHC_6H_5)_2 + H_2O \rightarrow HCI +$

+ $830_2 \mathrm{MHPO} (\mathrm{HHO_0H_0})_2$. The salt formation of (11) by action of elkali liquors and amnonia takes place wit but difficulty. On an acidification of these easily soluble salts the free (III) compounds separate so that the reaction mixture for the production of (III) can be directly separated from the alkali liquor. The compounds (IE) are colorless belies of bittor taste and cannot be hydrolized with alkali solutions; this can, however, be achieved by a heating with dilutes minoral acids under the formation of emides of the alkylsulfo acids and anilities of the phosphoric acids. They are mono--basic, rantor strong acids. Trianilidophosphasosulfenalkydes (IV) are obtained in good yields on a longer heating of the trichlorophosphezosulfonelkyls with excess aniline in beazene solution. The properties of the products (II-IV) are mentioned in the experimental part. There are 3 references, 3 of which are Soviet.

ASSOCIATION: Card 2/3

Dnepropetrovskiy metallurgicheskiy institut (Dnepropetrovsk Metallurgical Institute)

Anilides of Alkylsulfonamidophospharic Acids

SOY/79-28-6-34/65

SUBMITTED:

February 21, 1957

1. Anilines--Chemical reactions

Card 3/3

AUTHORS: 304/ 79-28-6-35/63 Kirsanov, A. V., Levchenko, Ye. S.

TITLE: Fluoranhydride Salts of the Arylsulfonamidophosphoric Acids

(Soli ftorangidridov arilsul'fonamidofosfornykh kislot)

PERIODICAL: Zhurnal obshchey khimii, 1958, Vol. 28, Nr 6, pp. 1589-1594

(USSR)

ABSTRACT: In continuation of their own previous papers on trichloro-

phosphazosulfonaryls and on products of their hydrolysis, the dichlomanhydrides of the arylsulfonamidophosphoric acids (Refs 1, 2) the authors were interested in synthesizing and investigating the fluorine containing analogues of the trichlorophosphazosulfonaryls and of the dichloroanhydrides of the arylsulfonamidophosphoric acids. The direct substitution of the chlorine in these compounds by fluorine, as for instance, with potassium fluoride, did not succeed. As is known chlorine is easily replaced by fluorine in the chloroanhydrides of various acids when potassium fluoride in aqueous solution is allowed to act on them. In the conversion of the trichlorophosphazosulfonaryls with a saturated solution of

potassium fluoride heat is produced, on which occasion be-Card 1/3 sides the substitution of chlorine by fluorine a partial

sov/ 79-28-6-35/63

Fluoranhydride Salts of the Arylsulfonamidophosphoric Acids

hydrolysis takes part and potassium salts of the difluoranhydrides of the arylsulfonamidophosphoric acids are formed according to the summary scheme 1. The same products are obtained in the conversion of the dichloroanhydrides of the same phosphoric acids with potassium fluoride according to scheme 2. As was to be expected the potassium salts of the dichloroanhydrides of the arylsulfonamidophosphoric acids, the potassium salts of the difluoranhydrides of the arylsulfonamidophosphoric acids and the dipotassium salts of the monofluoranhydrides of the same acids are formed depending on the reaction conditions. The structure of the potassium salts of the difluoranhydrides of the same acids is proved by their conversion to the esters of these acids by means of sodium methylate. It was shown that the potassium salts of the difluoranhydrides of the arylsulfonamidophosphoric acids have a still greater resistance to hydrolysis than the potassium salts of the dichloroanhydrides of the same acids. The reasons for this phenomenon are discussed in detail. There are 3 tables and 10 references, are Soviet.

Card 2/3

SOV /79-28-6-35/63

Fluoranhydride Salts of the Arylaulfonamidophosphoric Acids

ABSOCIATION: Institut organicheskoy khimii Akademii nauk Ukrainskoy SSR (Institute of Organic Chemistry, AS Ukr SSR)

February 21, 11957 SUBMITTED:

1. Phosphoric scid-livdrolysis

Card 3/3

CIA-RDP86-00513R000722720004-7" APPROVED FOR RELEASE: 06/13/2000

"APPROVED FOR RELEASE: 06/13/2000 CIA-RDP86-00513R000722720004-7

AUTHORS:

Kirsanov, A. V., Nekrasova, Z. D.

301 / 79-28-6-36/63

TITLE:

The Diphenylamide of Trichlorophosphazocarbonic Acid and Its Derivatives (Difenilamid trikhlorfosfazougol'noy kis-

loty i yego proizvodnyje)

PERIODICAL:

Zhurnal obshchey khimii, 1958, Vol. 28, Nr 6, pp. 1595-1601

(USSR)

ABSTRACT:

Only one method for the synthesis of the N-phosphoric acid derivatives of urea (carbamidephosphoric acids) has been published until now, that is to say, the binding of the published until now, that is to say, the binding of the primary and secondary amines to the chlorine anhydride or primary and secondary amines to the chlorine anhydride or to the isocyanatephosphates (Ref 1). Besides, a bis-tri-to the isocyanatephosphates (Ref 2) was obtained on the action chlorophosphazocarbonyl (Ref 2) was obtained on the action of phosphorpentachloride on urea; this product being a large transfer of the N.N'-carbamide biphosphoric acid. In order

or prosphorpentachloride on the sphorphoric acid. In order derivative of the N,N'-carbamide biphosphoric acid. In order to develop a common method of synthesis for the N,N-double substituted carbamide-N'-phosphoric acids and their derivatives, and at the same time to extend the possibilities for using the phosphorpentachloride reaction with acid amines, the reaction of phosphorpentachloride with N,N-diphenylurea was carried out. Phosphorpentachloride reacts on N,N-di-

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The state of the s